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- (A) Chemically modified lympholdne and production thereof.
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- Proprietor: Takede Chemical Industries, Ltd. 27, Doshomechi 2-chome Higashi-ku Qseka-chi Osaka, 541 (JP)
- (inventor: Mishimura, Caamu 123-502, 2 Higashitoyonakacho 5-choma Toyonaka Caaka 560 (JP) Inventor: Pujina, Masshiko 10-7, Higarigaaka 2-choma Takarazuka Hyogo 665 (JP)
- Representative: von Kreisler, Alek, Dipl.-Chemet et el Delchmannhaus am Hauptbehnhof D-6000 Köln 1 (DE)

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Description

Lymphokines such as interferons (hereinafter sometimes abbreviated as IFNs) and interleukin-2 (hereinafter sometimes abbreviated as IL-2) have been considered to be of clinical value for the treatment of virol infections and malignancies and recent technological advances in genetic angineering have made it in principle possible to produce such lymphokines on large scales. However, it is known that the clearance of lymphokines administered to the living body is in general very short. In the case of lymphokines derived from heterologous animals, it is enticipated that entibodies may be produced in some instances and cause asvere reaction such as anaphylaxis. Therefore, technology development is desired which leads to delayed clearance of lymphokines used as drugs, with their ectivity retained, and further to decrease in their antigenicity. To achieve this object, chemical modification of lymphokines is a very effective means. Such chemical modification is expected to result in delayed clearance in the living body, decreased antigenicity and, further, increased physiological activity. From the practical viewpoint, the significance of chemical modification of lymphokines is thus very great.

Generally, in chemically modified physiologically active proteins, a method is required by which said proteins can be chemically modified while retaining their physiological activity. Polyethylene glycol methyl ether is considered to have no entigenicity and therefore is used in chemical modification of proteins. The introduction of said substance into proteins is generally performed by way of the intermediary of cyanuric chloride. However, cyanuric chloride is toxic per se and the possible toxicity of its degradation products in vivo remains open to question. Therefore, cyanuric chloride should be used with caution. Furthermore, the reaction involved requires a pH on the alkaline side and therefore the above-mentioned method of modification has a drawback in that it cannot be applied to proteins liable to inactivation under alkaline

U.S. Patent No. 4,002,531 discloses a method of producing monoalkylpolyathylane glycol derivatives of enzymes. However, the method disclosed therein, which uses sodium borohydride at pH 8.5, when applied to lymphokines, may possibly destroy the physiological activity of lymphokines and therefore cannot serve as an effective method of production. Furthermore, said patent specification does not any suggestion as to the effect of dalaying the *in vivo* clearance of the enzyme derivatives. Such effect is therefore unknown.

There is also known a method of introducing a low molecular aldehyde such as formaldehyde, acetaldehyde, benzaldehyde or pyridoxal into physiologically active proteins in the presence of a boron-containing reducing agent (Methods in Enzymelogy, 47, 489—478 (1977); Japanese Patent Unexamined Publication No. 154,596/83]. However, application of said method to lymphoidines falls to achieve effective delay in clearance. A substantial decrease in antigenicity cannot be expected but rether it is possible that the low molecular aldehyde introduced may saive as a hapten to thereby provide said lymphoidines with Immunogenicity.

The present inventors studied intensively to overcome the above difficulties and have now completed the present invention.

This invention provides chemically modified tymphokines having polyathylene glycol of the formula

wherein R is a protective group for the terminal oxygen atom and n is an optional positive integer, bonded directly to at least one primary amino group of the lymphokine moiety and a method of producing the same.

In the present specification, the term "lymphokine" includes soluble factors released from lymphocytes and involved in cellular immunity and substances equivalent thereto in physiological activity.

Thus, the lymphokines may be genetically engineered products, products derived from various enimals including humans and further include substances similar in structure and in physiological activity to these.

For instance, there may be mentioned various interferons (interferon-a (IFN-a), interferon-\(\beta\) (IFN-a), interferon-\(\be

Examples of said substances similar in structure and in physiological scilvity are substances having the structure of IFN-y except for the lack of 2 to 4 amino acids at the N-terminal thereof (PCT/JP84/00292, filed June 6, 1984), various IFN-y fragments tacking in the C terminal portion of IFN-y (e.g. 15K species; EPC Petent Application No. 86 111133.9), substances having the structure of IL-2 except for the lack of the N-terminal amino acid thereof (EPC (laid open) 91539) or the lack of 4 amino acids from the N-terminal (Japanese Petent Application 58-235638, filed December 13, 1983) and substances having the structure of IL-2 except for the lack of one or more constituent amino acids with or without one or more substitute amino acids in place of said missing one or ones, for example the IL-2 analog containing serins in lieu of the 125th amino acid cysteine (EPC (Isld open) 104798).

Preferred arriging such lympholenes are IFN-q. IFN-y (consisting of 146 amino acids (EPC (laid open) 0089676)), IFN-y lacking in two N-terminal amino acids (IFN-y d2), IFN-y lacking in three N-terminal amino acids (IFN-y d3), and IL-2.

The tymphokines to be used in the practic of the invention preferably have a molecular weight of 5,000 to 50,000, more preferably 10,000 to 30,000.

The primary amino group of lymphokines includes the N-terminal d-amino group and the e-amino

group of the lysine residue.

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Referring to the group represented by the above formula (I), the terminal oxygen-protecting group R is, for example, an alkyl or elkanoyl group. The alkyl group is preferably an alkyl of 1 to 18 cerbon etoms, more preferably a lower (C₁₋₄) elkyl, such as methyl, ethyl, propyl, i-propyl, butyl, l-butyl, sec-butyl or t-butyl. The alkanoyl group is preferably an alkanoyl of 1 to 8 cerbon stoms, more preferably a lower (C₁₋₄) elkanoyl, such as formyl, acetyl, propionyl, butyryl, i-butyryl or caproyl. The positive integer n is preferably not more then 500, more preferably 7 to 120.

The group of formula (i) preferably has a molecular weight of not more than 25,000, more preferably 350 to 6,000. From the viewpoints of physiological activity retention and clearance delaying effect, the group of formula (i) preferably has a molecular weight corresponding to 1 to 10%, more preferably 2 to 6%

of the malecular weight of the lymphokine to be modified.

The chemically medified lympholines according to the invention have the group of formula (I) directly

bonded to at least one of the primary group of the corresponding lymphokines.

When the N-terminal q-amino group is the only primary amino group in the lymphokine to be modified, the modified lymphokine has the group of formula (I) directly bended to said amino group. When the lymphokine to be modified has one or more lysine residues in its molecule, the modified lymphokine has the group of formula (I) directly bonded to some percentage, preferably 15 to 80% (on the average), of said s-amino groups. In this case, the N-terminal q-amino group may have or may not have the group of formula (I) directly bonded thereto.

The chemically modified lympholines according to the invention can be produced, for example, by

reacting a lymphokine with the aldehyde of the formula

wherein R and n are as defined above, in the presence of a reducing agent.

As the boron-centaining reducing agent to be used as conducting the reaction, there may be mentioned sodium borohydride and sodium cyanoborohydride. Among them, more preferred is sodium cyanoborohydride from the viewpoint of selectivity of reaction or possibility of carrying out the reaction in the neighborhood of neutrality.

In carrying out the reaction, the aldehyde (II) is used in an amount of about 1 to 10,000 moles per mole of the tymphokine, and the boron-containing reducing agent is used in an amount of about 1 to 100 moles per mole of the lymphokine. The degree of modification can be selected as desired by varying the mole ratio between lymphokine and aldehyde (II). The solvent to be used in carrying out the invention may be any solvent which does not disturb the reaction and is, for example, a buffer such as a phosphate or borate puffer. An organic solvent which does not inactivate lymphokines or disturb the reaction, such as a lower alkenol (e.g. methanol, ethanol, i-propanol) or acetonitrile, may be added. The reaction may be conducted within a broad pH range of 3 to 14 but is preferably performed in the vicinity of neutrality (pH 6.5—7.5). The reaction temperature may be selected within a broad range of 0° to 80°C, preferably 0° to 50°C, so as not to esuee denaturation of lymphokines. A period of 0.5 to 100 hours, generally 10 to 80 hours, will be sufficient for the reaction. The desired, chamically modified lymphokines can be obtained by purifying the reaction mixture by distysis, salting out, ion suchange chromatography, gel filtration, high performance flould chromatography, electrophoresis, or the like ordinary method of purifying proteins. The degree of modification of the amino group or groups can be celculated by sold degredation followed by amino acid analysis, for instance.

The above-mentioned sidehyde (ii) can be produced from an ethylene glycol derivative of the formula

wherein R and n are so defined above, for instance. The following is a method of producing the same which is advantageous in that the production of the corresponding byproduct carboxytic acid is little.

Thus, the compound (III) is exidized with pyridinium chlorochromets in a helosikane solvent such as methylene chloride or chloroform. In this case, pyridinium chlorochromate is used in an amount of 1 to 3 moles per mole of compound (III) and the reaction is carried out at -10° to 50°C, preferably at room temperature, for 1 to 30 hours.

Treatment of compound (III) (n-1) with potassium butoxide in t-butanol followed by reaction with a bromoscatal and treatment with an acid such as an organic acid (e.g. trifluoroscatic acid) or an inorganic acid (e.g. hydrochloric or sulfuria acid) can also give the corresponding sidehyde (II) which is longer in chain langth by —0—CH₂CH₂— then compound (III). In this case, 10 to 30 males, per male of compound (III), of potassium t-butoxide is added to the above compound and, after dissolution, 3 to 15 males, per male of compound (III), of a bromoscatal is added, followed by reaction at 10° to 80°C for 0.5 to 5 hours. After treatment of the reaction mixture in the conventional manner, the product is dissolved in a dilute aqueous solution of the above-mentioned acid, followed by heating for 5 minutes to 2 hours.

in each case, the reaction mixture can be subjected to purification process conventional in the field of chemistry, such as extraction, concentration, recrystallization, reprecipitation, chromatography and/or distillation.

The chemically modified lymphokines according to the invention have useful physiological activities similar to those of the corresponding known, unmodified lymphokines and are useful as drugs, among others.

The chemically modified lymphokines according to the invention exhibit delay in clearance in vivo as compared with the corresponding known, unmodified lymphokines and are low in taxicity and antigenicity and can be used safely for the same purposes and in the same manner as in the case of known by lymphokines.

The chemically medified tymphokines according to the invention can usually be administered to mammals (monkey, dog, pig. rabbit, mouse, human) either orally or parenterally in the form of appropriate pharmaceutical compositions prepared by using carriers, diluents, etc., which are known in themselves. Thus, for instance, chemically modified IFN-a according to the invention, when used as an antiviral

Thus, for instance, chemically modified IFN-a according to the invention, when used as an antiviral egent, is recommendably administered to human adults once a day by intravenous injection in a dose of 1×10° to 1×10° international units.

In the present specification, the amino scids, when referred to by abbreviations, are abbreviated according to IUPAC-IUB (Commission of Biological Nomenclature).

The transforment Escherichia coli 294/pHTtrp1101-d2 as disclosed hereinleter in a reference example has been deposited with institute for Fermentation, Osaka (IFO) under the deposit number IFO-14360 and, since June 6, 1984, with the Fermentation Research Institute (FRI), Agency of Industrial Science and Technology, Ministry of International Trade and Industry under the deposit number FERM BP-703 under

Budspest Treaty.

The strain Escherichie coli DH1/pTF4 has been deposited with the Institute for Fermentation, Osaka under the deposit number IFO-14299 and, since April 6, 1984, with the FRI under the deposit number FERM BP-628 under Budspest Treaty.

Brief description of drawings

Fig. 1 shows the clearance-delaying effect in rat plasma as disclosed in Example 1 (iv). The measurement results obtained with the chemically modified IFN-α according to the invention as produced in Example 1 (i) are indicated by () (anxyme immunoassay) and () (antiviral activity assay), and the results obtained with riFN-αA used as a control by () (anzyme immunoassay) and () (antiviral activity assay). Fig. 2 shows the clearance-delaying affect in rat plasma as disclosed in Example 3 (ii). The data indicated by Δ, () and () are the enzyme immunoassay data for compound No. 8, compound No. 2 (Table

1) and control rFN-aA, respectively.

Fig. 3 shows the construction scheme for the expression plasmid pHITtrp1101-d2 disclosed in Reference Example 3 (i) and Fig. 4 the construction scheme for the expression plasmid pLC2 disclosed in Reference Example 4 (i).

Best mode for corrying out the invention

The following working examples and reference examples illustrate the invention in more detail but are by no means limitative of the invention.

Example 1

6 Production of polyethylene glycol methyl ether-modified IFN-a

(I) A 5-mi (4.8 mg as protein) portion of a solution of IFN-d (rIFN-dA) was dialyzed against 0.2 M phosphate buffer (pH 7.0) and 0.15 M apdium chipride at 4°C for 12 hours. To the dislyzate taken out, there was added the polyethylaneglycal methyl ether aldehyde (everage malecular weight 1,900) (260 mg) obtained in Reference Example 1. Then, sodium cyanoborohydrida (140 mg) was added, and the mixture was stirred at 37°C for 40 hours. The reaction mixture was poured into a Sophadex G-75 column (3.0×43.0 cm) and developed with 25 mM ammonium acetate buffer (pH 5.0) and 0.15 M sodium chloride. The cluste was collected in 8-mi portions. Eluste fractions (190-150 ml) containing the contemplated product were combined. Asserying by the Lowry method using bovine serum albumin as a standard revealed that the protein content in the combined fractions was 84 µg/ml. Amino acid ratios in acid hydrolysats (6 N hydrochloric acid, 110°C, 24 hours) were as follows: Asp, 12.2 (12); Thr, 10.4 (10); Ser, 16.0 (14); Giu, 24.8 (28); Pro, 6.0 (5); Gly, 6.2 (5); Ala, 8.6 (8); Val, 6.5 (7); Mar, 4.0 (5); Ila, 7.6 (8); Lau, 21.0 (21); Tyr, 6.2 (5); Pho, 9.9 (10); Lys. 6.5; His. 3.8 (3); Arg. 9.1 (9); Cys. Trp., decomposed. In view of the fact that HFN-QA contains 11 Lys residues, the above results led to a conclusion that about 41% of Lys residues in interferon a had been modified at the e-emino group with the polyethylane glycol methyl ether (sverage molecular weight 1,900). The potency of this product as determined by the enzyme immunoassay method [Methods in Enzymology, 79, 589—596 (1981)] was 1.51×107 international units/mg and the antiviral activity as determined by the method described in Journal of Virology, 37, 755—758 (1981) was 0.57×107 international units/mg. This product (IFA-3) was submitted to a clearance test in rate as mentioned later herein.

(ii) Using 100 mg of the polyethylene glycol methyl ether aldehyde obtained in Reference Example 1 and having an average molecular weight if 750 and 100 mg of sodium cyanoborohyddde, riFN-qA was

treated in the same manner a (i) to give 30 ml of a solution of polyathylane glycol methyl ether-modified IFN-a with a protein content of 130 µg/ml. Amino acid ratios in acid hydrohysate (6 N hydrochloric acid, 110°C, 24 hours) were as follows: Asp, 12.1 (12); Thr, 10.1 (10); Ser, 13.6 (14); Glu. 25.7 (26); Pro, 5.5 (6); Gly, 5.6 (5); Ala, 8.4 (8); Val, 6.7 (7); Mat, 5.5 (5); Ila, 7.4 (8); Leu, 21.0 (21); Tyr, 5.1 (5); Pho, 9.6 (10); Lya, 4.7; 6 His, 3.5 (3): Ara, 9.1 (9): Trp. 1.8 (2); Cys, decomposed. The above data indicate that about 57% of Lys residues had been modified at the s-amino group. Enzyme immunossesy performed in the same manner as (i) gave the result 5×10° international units/mg, and the antiviral activity of the product was 0.14×10° International units/mg.

(iii) The procedure of (i) was followed using 27 mg of the polyethylene glycol methyl ether aldehyde and 27 mg of sodium cyanoborohydrida and there was obtained 50 ml of a polyethylene glycol methyl other-modified IFN-a solution with a protein content of 45 µg/ml. Amino acid ratios in acid hydrolysate (8 N hydrochloric acid, 110°C, 24 hours) gave the following results: Asp, 13.8 (12); Thr, 10.4 (10); Ser. 14.9 (14); Glu, 28.6 (26); Pro. 5.5 (5); Gly. 6.1 (5); Ala, 8.3 (8); Val. 6.6 (7); Met. 5-2 (5); Ile, 7.4 (8); Leu, 21.0 (21); Tyr, 5.3 (5); Phe, 10.2 (10); Lye, 9.0; His, 3.8 (3); Arg, 9.1 (9); Trp, 2.3 (2); Cya, decomposed. The above data indicate that about 18% of Lys residues had been modified at the e-amino group. Enzyme immunoassay performed in the same manner as (1) gave the result 1.09×10⁸ international units/mg and the antiviral activity of this product was 1.53×10⁸ international units/mg.

(IV) The chemically modified IFN-d (IFA-3) of the invention as obtained above in (I) was administered to e group of three 7-week-old female SD rats by injection into the femoral muscle in a dose of 1.274×10⁴ units per capita. After a prescribed period, blood was sampled from the caudal vein and the IFN-a potency in plasma was determined by the enzyme immunosassy method and antiviral activity method described in Example 1 (I). A distinct dolay in clearance was observed as compared with a group administered unmodified interferon a (riFN-aA) in a dose 1.259×10° units per capita.

The above results are depicted in Fig. 1.

Example 2

To 5 mi of the solution of chemically modified IFN-c (IFA-3) of the invention as obtained in Example 1 (I), there is added 250 mg of human serum albumin. The resulting solution is flittered through a membrane filter (pore size: 0.2 µm) and distributed into 5 vials, followed by lyophilization and storage. The contents of each visi are dissolved in 1 ml of distilled water for injection just prior to use.

Example 3 Production of polyethylene glycol methyl ether-modified IFN-a end alkanoyl-polyethylene glycol-modified

(i) The tide compounds were synthesized by using the polyethylane glycol methyl ether sidehyde and alkaney/palyethylene glycol sidehyde obtained in Reference Example 1 and Reference Example 2, respectively, and following the procedure of Example 1. Various data for each derivative synthesized are shown in Table 1 and amino sold analysis data therefor in Table 2.

(II) The chemically modified IFN-a species obtained in (I) above (compounds No. 2 and No. 8) were administered to 7-week-old female SD retain groups of 3 by intramuscular injection into the femur in doses of 3.12×10° units and 2.66×10° units, respectively. Thereafter, blood samples were collected from the caudal vein at times intervals and assayed for IFN-a potency in plasma by anxyme immunoassay. Obviously delayed clearance was noted as compared with the group given 3.82×10° units of unmodified IFN-a. These results are depicted in Fig. 2.

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TABLE ?
Polyethylens glycal methyl athar-modified interferon a and attanoyl polyethylens glycal-modified interferon a

EIA	2,02×10 ⁷ 8.63×10 ⁹	1.30×10' 5.63×10°	6.00×10° 1.58×10°	3.31×10*	2.60×10 ⁷	4.70×107	1.28×10' 2.85×10'	1.77×10 ⁷	2.67×10' —
% Modi- fica: tien	. 31	8 2	3.6	2	9	916	20	3	99
Yield (%)	68	79	100	73	84	70	16	88	73
Ob- taines (ml)	36	22	92	17.5	98	97	38	92	3£
Content OD 280 am	0.139	0.151	0.210	a 175	0.100	0.117	0.107	0.160	0.087
Reac- tian time (hours)	18	16	10	18	24	48	78	24	24
NaBH ₅ CN smount (mg)	50 (cs. 200 times)	64 (ca. 200 times)	62 (cs. 200 times)	50 (ce. 200 times)	60 (ca. 200 times)	100 (ca. 400 times)	100 (cs. 400 times)	60 (ca. 240 timas)	50 (ca. 200 times)
Addition of NeBH,CN	Sarue time	Seme	Serne	3 hrs leter	5 hrs later	24 hrs later	5 hrs later	7.5 hrs fater	8 has fater
PEG eldehyde enount (mg)	252 (ca. 20 times)	124 (ca. 10 times)	61 (ca. 6 timent)	47 (ca. 10 times)	110 (oa. 60 tlmes)	56 (ca. 70 times)	162 (cs. 120 times)	184 (ca. 50 times)	120 (a. 60 dmes)
1 de de c	37	B	37	33	4	•	v. 🕶	•	₹.
PEG aldehyde (sv. mol.	MeOPEG (MeOPEG (5000)	MeOPEG (Septi)	MeDPEG (1908)	MeOPEG (750)	MeOPEG (550)	MeOPEG (350)	Acetyl PEG 1540)	Capitaly PEG (1100)
(FIV-ci assrount	5 ml (42 mg)	6 ml (4.2 mg)	5 ml (4.2 mg)	6 ml (4.2 mg)	6 ml (4.2 mg)	5 ml (4.2 mg)	5 ml (4.2 mg)	6 mf (4.2 mg)	5 ml (4.2 mg)
Com- pound No.		2	m	4	æ	6	7	60	6 5

PEG: Potyathidane giyes), MsOPEG: Potyathylene giyesi methyl ether, The vatue in perentheses is the evenge molecular weight. NaBHyCN: Sodium cyanobovohydride, ElA: Enzyme Immunosesey, AVA: Anihinel ectivity

TABLE 2

		aulav eleylana bise enimA										
s	Com- pound Na.	1	2	8	4	5	6	7	8	9	rIFN -cA	Theo- retical value
	Asp	12.8	12.7	12.5	12.5	13.4	12.9	12.2	12.5	12.8	12.6	12
10	Thr	11.7	11.6	11.2	10.9	11.3	11.4	10.9	11.6	11.3	11.6	10
	Ser	15.8	16.7	16.7	15.4	17.8	16.6	15.4	16.8	15.6	15.6	14
15	Glu	27.4	27.0	26.7	27.3	27.8	27.3	26.1	20.3	26.4	27.6	26
	Pro	_	5.3	5. 6	5.5	5.6	5.8	5.5	6.7	5.7	3.7	6
	Gly	4.9	5.0	4.6	4.6	7.1	4.8	4.5	5.3	5.4	4.6	5
20	Als	8.1	8.0	8.1	7.8	8.8	7.5	7.3	8.3	8.4	7.8	8
	Сув		_	_		_	<u> </u>	-	_	-	_	4
25	Val	6.8	6.8	6.7	6.6	7.3	6.7	6.3	6.9	7.1	6.6	7
}	Met	3.2	4.7	4.3	4.3	44	4.3	4.1	4.7	4.8	3.9	5
	lle	7.7	7.7	7.7	7.6	6.0	7.6	7.3	7.5	7.6	7.6	8
30	Leu	21.0	21:0	21.0	21.0	21.0	21.0	21.0	21.0	21.0	21.0	21
i	Тут	4.3	4.5	4.6	4.6	4.8	4.6	44	4.8	4.8	4.6	5
3 5	Phe	9.8	9.8	9.8	9.8	9.8	9.8	9.4	9.7	9.8	9.8	10
	Lys	8.6	10.3	10.5	9.6	5.4	6.1	2.3	6.6	4.9	11,3	11
	His	2.7	2.0	2.7	2.7	2.9	2.8	2.6	2.9	2.9	4.1	3
40	Arg	8.8	8.8	9.2	8.8	9.1	8.8	8.5	7.7	7.6	8.9	9
	Тпр	-	-	-	_	_	-	_	0.8	1.0	_	2

-: Not detected.

Example 4
Production of polyethylene glycol methyl ather-modified interferenty

(i) A 5-ml portion (5.95 mg as protein) of a solution of the interferent produced by the recombinant DNA technique (hereinafter abbreviated as riFN-y; cf. EPC laid open No. 110044) was applied to a Sephadex G-25 column (2.0×50.0 cm) and developed with 0.2 M phosphata buffer (pH 7.0). The eluste was fractionated in 5-ml portions. Fractiona Nos. 11—12 were combined and diluted to 100 ml with the same buffer. Thereto was added polyethylene glycol methyl ether aldahyda (average molecular weight 750) (225 mg), followed by addition of sodium cyanoborohydrida (300 mg). The mixture was shaken at 37°C for 72 hours. The resulting propipitate was removed by centrifugation. The supermatant was concentrated to 19 ml using a Claffow membrane (Amicon). The concentrate was applied to a Sephadex G-75 column (3.0×43.0 cm) and developed with 25 mM ammonium acetate buffer (pH 6.0)+0.15 M sodium chloride+10 mM glutathions. The cluste was fractionated in 5-ml portions. Fractions Nos. 17—24 containing the desired product were combined. The protein content in the combined fractions as determined by the Bradford method using bovins aerum albumin as a standard was 7.73 µg/ml. The acid hydrolysate (6 N hydrochloric scid, 110°C, 24 hours) gave the following amino acid analysis values: Asp, 19.8 (20); Thr, 4.7 (5); Sor, 8.3 (11), Glu, 18.5 (18); Pro, 2.1 (2); Gly, 5.4 (5); Ala, 7.8 (8); Val, 8.4 (8); Mat, 3.7 (4); Ila, 7.1 (7); Leu, 9.7 (10), Tyr, 5.3 (5); Phe, 9.7 (10); Lys, 17.6; His, 2.0 (2); Arg, 5.0 (8); Cys, Trp, decomposed. Since riFN-y contains 20 Lys realdues, the above results indicate that about 12% of the Lys s-amino groups in riFN-y on an artiviral activity

of 1.3×10⁴ International unite/mg. Administration of the product to rate resulted in obvious delay in clearance in blood. On the other hand, the precipitate was dissolved in 6 M guanidine hydrochloride and dialyzed against 25 mM ammonium scetate (pH 8.0)+0.15 M sodium chloride+10 mM glutathione at 4°C overnight, followed by Sephadex G-75 gel filtration in the same manner as above. The thus-purified fraction (25 ml) had a protein content of 128 µg/ml and amino acid enalysis of the acid hydrolysate (8 N hydrochloric acid, 110°C, 24 hours) gave the following values: Asp, 20.0 (20); Thr, 5.2 (5); Ser, 9.5 (11); Giu, 27.8 (18); Pro, 27 (2); Giy, 14.5 (5); Ala, 8.1 (8); Val, 8.5 (8); Met, 4.3 (4); ile. 7.2 (7); Leu, 10.2 (10); Tyr. 5.8 (5); Phe, 10.1 (10); Lya, 14.7; His, 2.0 (2); Arg, 7.3 (8); Thr, 0.7 (1); Cys, decomposed. The higher values for Giu and Giy than the theoretical are presumably due to contamination by glutathione. Since rIFN-y contains 20 Lya e-amino groups, the above results indicate that about 26.5% of the Lya e-amino groups in rIFN-y had been modified by polyethylene glycol methyl ether.

(ii) Using 225 mg of polyethylane glycol methyl ether aldehyde having an average molecular weight of 750 and 120 mg of sodium cyanoborohydride, riFN-v was treated in the same manner as (i) in the presence of 2-mercaptoethanol (2%) to give 30 ml of a polyethylane glycol methyl ether-modified riFN-y solution having a protein content of 236 µg/ml. Amino acid analysis of the acid hydrolysate (8 N hydrochloric acid, 110°C, 24 hours) gave the following values: Asp, 20.0 (20); Thr. 5.2 (5); Ser. 9.6 (11); Glu. 33.6 (18); Pro. 1.8 (2); Gly, 19.9 (5); Ala, 9.2 (8); Vei, 8.9 (8); Met, 4.8 (4); Ile, 7.4 (7); Leu, 10.2 (10); Tyr. 5.8 (5); Pho, 10.7 (10); Lys, 10.2; His, 2.3 (2); Arg, 7.9 (8); Trp. 0.6 (1); Cys. decomposed. The higher values for Glu and Gly are presumebly due to contamination with glutathions. Since riFN-y contains 20 Lys s-amino groups, the above results indicate that about 50% of the Lys s-amino groups in riFN-y had been modified by polyethylene glycol methyl ether.

Example 5

Production of polyethylene glycal methyl ether-madified IFN-yd2

(i) A 5-ml portion (4.85 mg as protein) of the IFN-yd2 solution obtained in Reference Example 3 is applied to a Sephadex G-25 column (2.0×50.0 cm) and developed with 0.2 M phosphate buffer (pH 7.0). The cluste is fractionated by 5 ml. Fractiona Nos. 11-13 are combined and diluted to 100 ml with the same buffer. To the dilution is added polyethylene glycal methyl other aldehyde (everage molecular weight 750) (200 mg), and then sodium cyangborohydride (300 mg). The mixture is shaken at 37°C for 72 hours. The resulting precipitate is removed by centrifugation. The supernature is concentrated to 10 ml using a Diaffirm membrane (Amicon). The concentrate is applied to a Sephadex G-75 column (3.0×43.0 cm) and developed with 25 mM emmanium acetate buffer (pH 0.0)+0.15 M sodium chloride+10 mM glutathione. The cluste is fractionated by 5 ml, and the fractions containing modified IFN-yd2 having the polyethylene glycol methyl ether moisty on the Lys s-amino group in the molecula are collected and combined. When this product is administered to rote, evident dalay in clearance in blood is noted.

On the other hand, the pracipitate is dissolved in 6 M guanidine hydrochloride, dislyzed against 25 mM ammonium acetate buffer (pH 6.0)+0.15 M sodium chloride+10 mM glutathione at 4°C overnight, and purified by Sephadex G-75 gel filtration in the same manner as above. Thus is obtained a fraction containing modified iFN-yd2 having the polyethylene glycol methyl ethyl moiety on the Lys s-amino group

In the molecule.

Example 6

Production of polyothylene glycol methyl ether-modified IFN-y3

(i) A 5-mi (5.5 mg as protein) portion of the IFN-yd3 solution obtained in Reference Example 4 is applied to a Sephadex G-25 column (2.0×60.0 cm), followed by development with 0.2 M phosphate buffer (pH 7.0). The stuate is fractionated in 5-mt portions. Fractions Nos. 11—13 are combined, and thereto are added polyethylene givcol methylether aldehyde (average motecular weight 750) (225 mg) and then sodium cyanoborohydride (120 mg). The mixture is shaken at 37°C for 24 hours. The reaction mixture is applied to a Sephadex G-75 column (3.0×43.0 cm), followed by development with 25 mM ammonium accetate buffer (pH 8.0). This is obtained a fraction containing modified IFN-yd3 with the polyethylene givcol methyl ether moiety on the Lys s-amino group in the molecule. When this product is administered to rate, obvious delay in clearance in blood is observed.

Example 7

55 Production of polyethylene glycci methyl ether-modified IL-2

(i) A 5-ml (5.0 mg as protein) portion of the interlaukin 2 (hereinafter abbreviated as rit-2) obtained in Reference Example 5 was dislyzed against 0.2 M phosphate buffer (pH 7.15) for 12 hours. To the dislyzete was added polyethylane glycol methyl ather aldehyde (average molecular weight 750) (97 mg), and then sodium cyanoborohydride (100 mg). The mixture was attred at 37°C for 24 hours. The resultant precipitate was removed by centrifugation. The supernetant was dislyzed againt 6 mM ammonium accusts buffer (pH 5.0) for 5 hours. The dislyzete was applied to a Sephadex G-75 column (3.0×43.0 cm) and developed with the same solvent system. The cluste was fractionated in 6-ml portions. The desired product-containing fractions Nos. 21—29 were combined. The combined fraction had a protein content of 25 µg/ml as determined by the Bradford method using bovine serum albumin as a standard. The acid hydrolysate (8 N hydrochloric acid, 110°C, 24 hours) gave the following amino acid snelysis values: Asp, 12.0 (12); Thr, 12.5

(13); Ser, 7.1 (8); Gly, 18.6 (18); Pro, 5.5 (5); Gly, 2.2 (2); Al., 5.0 (5); Val, 3.7 (4); Met, 3.9 (4); Ile, 8.1 (8); Leu, 22.2 (22); Tyr, 3.0 (3); Phe, 8.0 (6); Lys, 7.3; Hls, 3.0 (3); Arg, 3.9 (4); Cys, Trp, decomposed. Since ril-2 contains 11 Lye residues, the above results indicate that about 23.6% of the Lys e-amino groups had been modified by polyethylene glycol methyl other. The IL-2 activity of the product as determined by the method of Hinuma et al. (Biochemical and Biophysical Research Communications, 709, 383—369 (1982)] which measures the growth of an IL-2-dependent mouse natural killer cell line (NKC3) with the [⁸H]-thymidine uptake into DNA as an index was 22,988 units/mg. When rill-2 is supposed to have an activity of 40,000 units/mg, the product is estimated to retain 57.7% of the activity. After administration of this product, obvious delay in clearance in blood was noted.

Reference Example 1

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Synthesis of polyethylene glycol methylether aldehyde

(I) Polyathylane giycol matnyl ether (5 g; average molecular weight 5,000) was dissolved in methylane chloride (100 ml) and then pyridinium chlorochromate (330 mg) was added. The mixture was stirred at room temperature for 12 hours. The maction mixture was diluted two-fold with methylane chloride and poured into a Florisii column (6×10 cm), and the column was weeked with methylane chloride and then with chloroform, followed by elution with methanolchloroform (1:8). Fractione positive to 2,4-dinitro-phenylhydrazine test were combined, the solvent was distilled off under reduced pressure, and there was obtained a crystalline wax. Yield 1.5 g (30%). Thin layer chromatography: R,=0.08 (chloroform-methanol-acetic acid=9:1:0.5, silice gel). ¹³C-NMR spectromatry revealed an absorption due to the aldehyde group in hydrated form [—GH(OH)] at 96.2 ppm.

(ii) Polyethylene glycol methyl ether (10 g; average molecular weight 5,000) was dissolved in tertiary-butenol (100 mi). Thereto was added patesium tertiary-butenolde (4.17 g), followed by addition of bromoecetal (2.56 ml). The mixture was stirred at 40°C for 2 hours. The tertiary-butanol was then distilled off under reduced pressure, water was added to the residue, and the equeous mixture was extracted with chloroform (200 ml x2). The extract was washed with water and dried over anhydrous sodium sulfate. The chloroform was then distilled off under reduced pressure, petroleum benzine was added to the residue, and the resultant cryatelline residue was collected by filtration and washed with other. Thus was obtained 9.5 g (95%) of the corresponding polyethylane glycol methyl ether disthyl scetal. A 5-g portion of the acetal was dissolved in 50 ml of 0.05 M trifluoroacetic acid, treated in a boiling water bath for 30 minutes and then hypothilized, giving a polyethylane glycol methyl ether aldehyde longer in chain length by —O—CH₂CH₂—than the product obtained in (i).

(iii) Polyethylene glycol methyl ether (5.7 g; everage molecular weight 1,900) was dissolved in methylene chloride (100 ml) and then pyridinium chlorochromate (870 mg) was added. The mixture was stirred at room temperature for 12 hours, then diluted with an equal volume of methylene chloride, and poured into a Florisli column (6.0×10.0 cm). The column was weshed with methylene chloride and then with chloroform. followed by clution with 10% methanol/chloroform Fractions positive to 2,4-dinitrophenylhydrozino test were combined. Removal of the solvent by distillation gave a crystalline wax. Yield 1.8 g (30%). Thin layer chromatography: R₂=0.10 (chloroform-methanol-acotic acid=8:1:0.5, silica gel).

**C-NMR spectrometry indicated the presence of an absorption due to the aldehyde group in hydrated form (—CH(OH)₂) at 98.2 ppm.

(iv) Polyethylena glycol methyl ethar (19.5 g; everage molecular weight 1,800) was dissolved in tertiary-butanol (100 ml). Potessium tertiary-butaxids (10.4 g) was added and then bromoscatel (6.4 ml) was added. The mixture was attred at 40°C for 2 hours. The tertiary-butanol was then distilled off under reduced pressure. Water was added to the residue, followed by extraction with chloroform (200 ml×2). The extract was washed with water and dried over anhydrous addition sulfate. The chloroform was distilled off under reduced pressure, petrolsum benzine was added to the razidue, and the resultant crystalline residue was collected by filtration and washed with ether to give 8.5 g (89.6%) of acetal. A 3-g portion of the scatal was dissolved in 0.08 M trifluoreacetic acid, and the solution was treated in a boiling water bath for 30 minutes and then tyophilized to give a polyethylene glycol methyl ether aldehyde langer in chain length by —O—CH₂CH₂— then the product obtained in (iii).

(v) Polyathylane glycol methyl ether species having everage molecular weights of 750, 550 and 350 were derived to the corresponding aldehyde species by following the above procedures,

55 Reference Example 2.

Synthesis of alkanovi polyethyleneglycol aldehyde

(i) In 50 mi of pyridine, there was dissolved 15 g of polyethylene glycol 1540 (Wake Pure Chemical Industries) (average molecular weight 1500). To the solution was added 1.85 mi of acetic anhydride. The mixture was stirred at 40°C for 2 hours and then at room temperature for 16 hours. Thereafter, the solvent was distilled off under reduced pressure. The residue was dissolved in chloroform, and the solution was distilled off under reduced pressure. The residue was dissolved in a small amount of chloroform, a petroleum benzine-ether (2:1) mixture was added to the solution, and the mixture was allowed to stand to give 14 g (90%) of a crystalline wax. A 1.4-g portion of the wax was dissolved in 50 mi of methylene chloride, followed by addition of 300 mg of pyridinium chlorochromate. The resulting mixture was stirred

at room temperature for 18 hours. The reaction mixture was applied to silice gel C-200 (Weko Puro Chemical Industries) column (3×50 cm), and the column was washed with 5% methanol-chloroform (200 ml) and eluted with 10% methanol-chloroform. Fractions positive to the 2.4-dinitrophenylhydrazine test were combined, and the solvent was distilled off under reduced pressure. A crystalline wax was obtained.

5 Yield 580 mg (41%).

(iii) In 50 ml of methylene chloride, there was dissolved 20 g of polyathylene glycol 1000 (Wake Pure Chemical Ind.) (average molecular weight 1000), followed by addition of 5.15 g of n-caproyl anhydride. The mixture was attred at 70°C for 2 hours. Then, the solvent was distilled off, and the residue was purified using a silica gel C-200 column (3×50 cm) and slutdon with ethyl acette-methanol (4:1) to give 14.9 g (60%) of an oil, which solidified upon standing in a refrigerator. The subsequent exidetion with pyridinium chlorochromate as conducted in the same manner as (i) gave the corresponding eldehyde.

Reference Example 3 — Production of IFN-yd2

(I) Transformant preparation

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The IFN-y expression plasmid pHITtrp1101 [cf. EPC (laid open) No. 110044, Example 2 (iii)) was digested with the restriction enzymes Avail and Pstl. and an Avail-Pstl 1 to DNA fragment containing the IFN-y gene portion was isolated. The protein synthesis start coden-containing oligonucleotide edapter

CGATAATGTGCCAG

TATTACACGGTCCTG

chemically synthesized by the phosphotniester method was joined to the above DNA fragment at the Avail cohesive and thereof using T4 DNA ligase.

The above adapter-joined gene was inserted into the DNA fragment obtained by cleavage of the plasmid ptrp771 [cf. above-cited publication, Example 2 (ii)] with the restriction enzymes C/el and Patl, downstream from the up promoter in said fragment. Thus was constructed the expression plasmid pHITtrp1101-d2 coding for the Cye-Tyr-deficient IFN-y polypoptide (Fig. 3).

Escherichie coll 284 was transformed with this plasmid pHftrp1101-d2 by the method of Cohen et al. (Proc. Natl. Acad. Sci. U.S.A., 69, 2110 (1972)) to give the transformant Escherichie coll (=E. call) 284/pHftrp1101-d2 carrying said plasmid.

(ii) Transforment cultivation

The strain E. coll 294/pH/Trp1101-d2 carrying the plasmid constructed in (i) above was cultivated in M9 medium containing 8 µg/mi of tetracycline, 0.4% of casemino acids and 1% of glucose at 37°C. When the growth reached KU 220, 3-p-indolylacrylic acid (IAA) was added to a concentration of 25 µg/mi. Thereafter, the cultivation was continued for further 4 hours. After cultivation, cells were horvested by contribugation and suspended in 1/10 volume of 0.05 M Tris-HCl (pH 7.6) containing 10% sucrose. To the suspension, there were added phenylmethylaulfonyl fluorido, NaCl, ethylenadizminetatracettate (EDTA), apermidine and lysozyme to concentrations of 1 mM, 10 mM, 40 mM and 200 µg/mi, respectively. After standing at 0°C for 1 hour, the suspension was treated at 37°C for 3 minutes to give a lysate.

The lysste was subjected to centrifugation at 4°C and 20,000 rpm (Serval) centrifuge, SS-34 rotor) for 30 minutes to give an IFN-yd2 polypeptide-containing suparasent. This supernatant had an entiviral activity of 2.97×10° Uniter culture fluid.

(iii) Purification of IFN-yd2

in 18 ml of 0.1 M Tris-hydrochloride buffer (pH 7.0) containing 7 M guanidine hydrochloride and 2 mM phenylmothylsulfonyl fluoride, there were suspended 5.9 g of cells obtained in the same manner as (ii) above and stored in the frozen state. The suspension was stirred at 4°C for 1 hour and then subjected to centrifugation at 10,000×g for 30 minutes to give 20 ml of a supernaturt. This supernaturt was diluted with 260 ml of a buffer (pH 7.4) comprising 137 mM sodium chloride, 2.7 mM potassium chloride, 8.1 mM disodium phosphate and 1.5 mM monopotassium phosphate (hereinafter such buffer being referred to by the abbreviation PBS) and the dilution was applied to an antibody column (Moy2-11.1, column volume 12 ml) at a flow rate of 1 ml/minute. The column was then washed with 00 ml of 20 mM sodium phosphate buffer (pH 7.0) containing 0.5 M guanidine hydrochloride and cluted with 38 ml of 20 mM sodium phosphate buffer (pH 7.0) containing 2 M guanidine hydrochloride to give 20 ml of an antivirally active fraction.

This 20-mi fraction was applied to a Sephacryl S-200 (Pharmacia) column (2.8×94 cm, column volume 600 ml) equilibrated in advance with 25 mM ammonium acetate buffer (pH 8.0) containing 1 mM ethylenediaminetetreacetate, 0.15 M aedium chloride, 10 mM cysteine and 2 M queniding hydrochloride, followed by ejution with the same buffer. Thus was obtained 37 ml of an antivirally active fraction.

The Cys-Tyr-deficient IFN-y polypeptide (IFN-yd2) obtained weighed 5.3 mg and had a pecific activity of 1.0×10° Ll/mg.

Reference Example 4 — Production of IFN-yd9

(1) Transforment production

The IFN-y expression plasmid pRC23/IFI-800 (cf. Example 7 of the specification for a patent application under EPC as laid open under No. 0089678] was digested with the restriction enzymes Ndel and Ncol, and a 710 bp Ndel-Ncol DNA fragment (A) containing the IFN-y gene region was isolated. Separately, the plasmid pRC23 was digested with the restriction enzyme Bg/II and EcoRI, and a 265 bp DNA fragment (8) containing the AP, promoter was isolated. The fregments (A) and (B) and the chemically synthesized, protein synthesis start codon-containing oligonucleotide

AATTCATGCAGGATCCA

GTACGTCCTAGGTAT

were joined together using T4 DNA ligase, with the Ndal and EmRi cohesive ends as the sites of joining. The DNA fregment thus obtained was joined to the plasmid pRC23/FI-600 after treatment with Neel and Bg/II, to thereby construct an expression plasmid, pLC2, coding for the Cys-Tyr-Cys-delicient IFN-y polypeptide (Fig. 2). This plasmid pLC2 was used for transforming Escherichia coli RRI(pRK248 cits) by the method of Cohen et al. (supra) to give a transformant, Escherichie coli)=E. coli) PRI(pLC2,pRK248 clts).

20 (ii) Transforment cultivation

The strein E. coli RRI(pLC2,pRK248 cits) carrying the plasmid constructed in (i) above was shake-cultured at 35°C in 60 ml of a liquid medium containing 1% Sectotryptone. 0.5% yeast extract, 0.5% sodium chloride and 7 µg/ml tetracycline. The culture broth was transferred to 2.5 liters of M9 medium containing 0.5% casemine sold, 0.5% glucose and 7 µg/ml tetracycline, and grown at 35°C for 4 hours and as then at 43°C for 3 hours. Cells were harvested by contritugation and stored at -80°C.

(III) Purification

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in 22 ml of 0.1 M Tris-hydrochloride buffer (pH 7.0) containing 7 M guanidine hydrochloride and 2 mM phenylmethylaulfonyl fluoride, there were suspended 7.1 g of frozen cells obtained in the same manner as mentioned above in (ii). The suspension was stirred at 4°C for 1 hour and then centrifuged at 10,000×g for 30 minutes to give 24 ml of a supernatant. This supernatant was diluted by adding 300 ml of PBS and the dilution was applied to an antibody column (Moy2-11.1, column capacity 15 ml) at a flow rate of 1 mil/minute. Theresiter, the column was washed with 60 ml of 20 mM sodium phosphate buffer (pH 7.0) containing 0.5 M guantdine hydrochloride and then sluted with 45 ml of 20 mM sodium phosphate buffer (pH 7.0) containing 2 M guantdine hydrochloride, to give 25 ml of an antivirally active fraction. This fraction (25 ml) was applied to a Sephacryl S-200 (Pharmacia) column (2.6×84 cm; column capacity 500 ml) aquilibrated in advance with 25 mM ammonium acetate buffer (pH 8.0) containing 1 mM athylenediamine tetrescetic sold, 0.15 M sodium chloride, 10 mM cysteine and 2 M guanidine hydrochloride, and sluted with the same buffer to give 40 ml pf an antivirally active fraction.

The thus-obtained Cys-Tyr-Cys-deficient IFN-y polypeptide IFN-y d3 weighed 7.0 mg and had a specific

activity of 2.72×107 IU/mg.

Reference Example 5 - Production of non-glycosylated human IL-2

(i) Transformant cultivation

E. coli DH1/pTF4 (EPC Pat. Appin. No. 84308153.0) was inoculated into 50 ml of a liquid medium (pH 7.0) containing 1% Bacto tryptone (Difco Laboratories, USA). 0.5% Bacto yeast extract (Difco Laboratories, USA), 0.5% sodium chloride and 7 µg/ml tetracycline as placed in a 250-ml Erlenmayer flask. After incubation at 37°C avernight on a swing rotor, the culture medium was transferred to a 5-liter jar fermenter containing 2.5 liters of M9 medium containing 0.5% casamino ecid, 0.5% glucose and 7 ug/mi tetracycline. Incubation was then conducted with sention and stirring at 37°C for 4 hours and after addition of 8-B-Indelylacrylic sold (25 µg/ml), for further 4 hours. Cells were harvested from the thus-obtained 2.5-liter culture broth by centifugation, frozen at -80°C and stored.

(ii) Extraction

The freeze-stored cells (12.1 g) obtained above were suspended uniformly in 100 ml of an extractant (pH 7.0) containing 7 M guanidine hydrochloride and 0.1 M Tris - HCl, the suspension was stirred at 4°C for hour and the lysate was centrifuged at 28,000×g for 20 minutes. There was obtained 93 ml of a supernetent.

(III) Purification of IL-2 protein

The supernatant obtained above was dislyzed against 0.01 M Trie HCI buffer (pH 8.5) and then centrifuged at 19,000×g for 10 minutes, giving 84 ml of a dislyzate supernatent. This dislyzate supernatent was applied to a DE 52 (DEAE-celluloss, Whatman, Great Britain) column (50 ml in volume) equilibrated with 0.01 M Tris · HCl buffer (pH 8.5) for protein adsorption. IL-2 was cluted making a linear NaCl concentration gradi nt (0—0.15 M NaCl, 1 liter). The active fractions (53 ml) were concentrated to 4.8 ml

using a YM-5 membrane (Amico, USA) and subjected to gel filtration using a Sephecryl S-200 (Pharmacia, Sweden) column (500 ml in volume) equibrated with 0.1 M Tris - HCI (pH 8.0)—1 M NaCl buffer. The active fractions (28 ml) obtained were concentrated to 2.5 ml using a YM-5 membrane. The concentrate was applied to an Ultrapore RPSC (Altax, USA) column for adsorption, and high performance liquid chromatography was performed using a trilluproacetic acid-acetonitrile system as the eluent.

Under the conditions: column, Ultrapore RPSC (4.8×75 mm); column temperature, 30°C; eluent A, 0.1% trifluoroscetic scid-99.9% water; sluent 8, 0.1% trifluoroscetic scid-99.9% acetonitriic; clution program, minute 0 (69% A+32% B)-minute 25 (55% A+45% B)-minute 35 (45% A+55% B)-minute 45 (30% A+70% B) - minute 48 (100% B); slution rate, 0.8 ml/min.; detection wave length, 230 nm, An active 10 fraction was collected at a retention time of about 39 minutes. Thus was obtained 10 ml of a solution containing 0.53 mg of non-glycosylated human IL-2 protein (specific activity, 40,000 U/mg; activity recovery from starting material, 30.6%; purity of protein, 99% (determined by densitometry)].

Claims

15

1. A chemically modified lymphokine having polyethylene glycol of the formula:

R+O-CH.CH.-L

wherein R is a protective group for the terminal oxygen atom and n is an optional positive imager, bonded directly to at least one primary amino group of the lymphokine mojety.

2. The modified lymphokine according to claim 1, wherein the lymphokine moiety has molecular weight from 5,000 to 60,000.

3. The modified lymphokine according to claim 2, wherein the lymphokine mojety has molecular 25 weight from 10,000 to 30,000.

4. The modified lymphokine according to claim 1, wherein the lymphokine molety is interferons, interlaukin-2, macrophage differentiating factor, macrophage activating factor, or substances similar in structure and in physiological activity to these.

6. The modified lympholine according to claim 1, wherein the lympholine moiety is interferon-a, interferon-β, Interferon-γ, interferon-γd2, interferon-γd3 or interleutin-2.

The modified lymphokine according to claim 1, wherein the lymphokine moiety is interferon-a.

7. The modified lymphokine according to claim 1, wherein the lymphokine moiety is interferon-y.

The modified lymphobine according to claim 1, wherein the lymphobine moisty is interleukin-2. 9. The modified lymphokine according to claim 1, wherein the polyothylane glycol has molecular

36 weight corresponding to 1 to 10% of the molecular weight of the lymphakine molecy. 10. The modified lymphokine according to claim 1, wherein the pulyethylene glycol has molecular

weight from 350 to 6,000.

11. The modified lymphokine according to claim 1, wherein R is alkyl or alkanoyl.

12. The modified lymphokine according to stelm 1, wherein n is a positive integer from 7 to 120. 13. The modified lympholine according to claim 1, wherein the primary amino group is N-terminal d-amino group or e-amino group of lysine residue in the lymphokine malety.

14. The modified lymphoidne according to claim 1, which has polyethylene glycol bonded to 15 to 80% of e-amino groups of tysine residue in the lympholone moiety,

15. A method of producing a chemically modified lymphokine having polyethylene glycol of the formula:

R-4-OCH_CH_-}

wherein R is a protective group for the terminal oxygen atom end n is an optional positive integer, bonded directly to at least one primary amino group of the lympholine moeity, which comprises reacting a lymphokine with an aldehyde of the formula:

R (O-CH, CH, -) -- CH, CHO

wherein R and n are as defined above, in the presence of a reducing agent.

16. The method according to claim 15, wherein the reaction is conducted in the neighborhood of neutrality.

17. The method according to claim 15, wherein the reducing agent is sodium cyanoborohydride.

Patantansprüche

1. Chemisch modifiziertes Lymphokin, das ein Polyäthylengiycol der Formal

R-I-O-CH.CH.--

worln R ine Schutzgruppe für das endständige Sauerstoffstom ist und nielne wählbere positive genze Zehl dersteilt, direkt an wenigstene eine primäre Aminogruppe des Lymphokinanteils gebunden enthält.

2. Modifiziertes Lymphokin nach Anspruch 1, worin der Lymphokinantell ein Molekulargewicht von 5.000 bis 50,000 besitzt.

3. Madifiziertes Lymphakin nach Anspruch 2. worin der Lymphokinanteil ein Moleculargewicht von 10.000 bis 30.000 sufwelst.

4. Modifiziertes Lymphakin nach Anspruch 1, worln der Lymphokinanteil aus Interferonen, interleukin-Makrophag-Differenzierungsfektor, Makrophag-Aktivlerungsfaktor oder diesen in Struktur und physiologischer Aktivität ähnlichen Substanzen besteht.

5. Modifiziertes Lymphokin nach Anspruch 1, werin der Lymphokinantail Interferon-a, Interferon-B;

Interferon-y, Interferon-ydz, Interferon-ydd oder Interieukin-2 ist.

6. Modifiziertes Lymphokin nech Anspruch 1, worin der Lymphokinantali Interferen-a ist.
7. Modifiziertes Lymphokin nech Anspruch 1, worin der Lymphokinenteli Interferen-y ist.

8. Modifiziertes Lymphokin nach Anspruch 1, worin der Lymphokinentell Interleukin-2 ist.

8. Mofiziertes Lymphokin nech Anspruch 1, worin des Polyäthylongiycol ein Maisulargewicht eufweist, des 1 bis 10% des Moleulargewichtes des Lymphokinanteiles entspricht. 15

10. Modifiziertes Lymphokin nach Anspruch 1, worin das Polyäthylangiyool ein Moleculargewicht von 350 bis 6.000 bositet.

11. Modifizierzes Lymphokin nach Anspruch 1. worin R für Alkyl oder Alkanoyl steht.

12. Modifiziertes Lymphokin nach Anspruch 1, worin n eine positive ganze Zahl von 7 ble 120 bedeutet. 13. Modifiziertes Lymphokin nach Anspruch 1, worin die primäre Aminogruppe eine N-endständige

a-Aminogruppe oder e-Aminogruppe eines Lysinrestes im Lymphokinantell derstellt.

14. Modifiziertes Lymphokin nach Anspruch 1, das ein Polyathylengiyool enthält, des en 15 bie 80% der a-Aminogruppen des Lysinrestes im Lymphokinantell gebunden ist.

15. Verfahren zur Herstellung eines chemisch modifizierten Lymphokins, das ein Polysthylenglycol der Formel

R-(-OCH_CH_-IR

worin R eine Schutzgruppe für das endständige Sauerstoffatom ist und n für eine wählbare positive genze Zahl steht, direkt an Wenigstens eine primäre Aminogruppe des Lymphokinanteils gebunden enthält, welches Verfahren die Umsetzung eines Lymphokins mit einem Aldehyd der Formel

R + O - CH_CH_ }_TO - CH_CHO,

worin R und n die vorstehend angeführte Bedeutung basitzen, in Gegenwert eines Reduktionsmittels umfaßt

16. Verfahren nach Anspruch 15, worin die Residion in der Nähe des Neutralbereiches durchgeführt wird.

17. Verfahren nach Anspruch 15, worin des Reduktionsmittel Natriumcyanborhydrid ist.

Revendications

35

1. Lymphokine chimiquement modifiée syant du polyéthylènegiycol de formule:

R-4-O-CH_CH_L

dans laquelle R est un groupe protectour de l'atome d'oxygène terminal et n cet un nombre antier positif laigaé su choix, lié directement à su moins un groupe amino primaire du fragment lymphokine.

2. Lymphokine modifiée solon la revendication 1. dans laquelle le fragment lymphokine a une masse moléculaire comprise entre 5000 et 50 000.

3. Lymphokine modifice selon la revendication 2, dans laquelle le fragment lymphokine a une masse motéculaire comprise entre 10 000 et 30 000.

4. Lymphokine modifiée selon le revendication 1, dans lequelle le fregment lymphokine ast un Interféren, l'Interleukine-2, un facteur de différenciation de macrophage, un facteur d'activation de macrophage, ou une substance similaire en structure et en activité physiologique è cas substances.

5. Lymphokine modifiée selon la revendication 1, dans laquelle le fragment lymphokine est l'interféron-a, l'interféron-B, l'interféron-y, l'interféron-yd2, l'interféron-yd3 ou l'interfeukine-2.

6. Lymphokine modifiés selon la revendication 1, dans lequelle le fragment lymphokine cat l'interféren-a

7. Lymphokine modifiée selon la revendication 1, dans lequelle le fragment lymphokine est l'interféron-y.

8. Lymphokine modifiée selon la revendicati n 1, dans laquelle le fregment lymphokine est l'Interleukine-2.

9. Lymphakine modifiée selon la revendication 1, dans laquelle le polyéthylènegiycol a une masse moléculaire correspondent à 1% à 10% de la masse moléculaire du fragment lymphakine.

10. Lymphokine modifiée salon la revendication 1, dans laquelle la polyéthylènegiyosi a une messe moléculaire comprise entre 350 et 6 000.

11. Lymphokine modifiés selon la revendication 1, dans lequelle R est un alkyle ou un alcanoyle.

12. Lymphokine modifiée selon la revendication 1, dans lequelle n'est un entier positif compris entre 7 et 120.

13. Lymphokine modifiée selon la revendication 1, dans laquelle le groupe amino primaire cet le groupe q-amino de l'extrémité N-terminale ou le groupe c-amino d'un reste lysine dans le fragment

10 lymphokine.

14. Lymphokine modifiée selon la revendication 1, qui a du polyéthylèneglycol lié à 15% à 80% des groupes s-emino du reste hysine dans le fragment lymphokine.

15. Procédé de préparation d'une lymphokine chimiquement modifiée ayant du polyéthylèneglycol de

R+O CH_CH_1

dans laquelle R est un groupe protectaur de l'atomo d'oxygène terminal et n est un nombre entier positif lalasé au choix. Ilé directement à su moins un groupe amine primaire du fragment lymphokine, qui comprend le réaction d'une lymphokine avec un aldéhyde de formule:

R-I-O--CH_CH_---O--CH_CHO

dans laquelle R et n sont tels que définis ci-dessus, en présence d'un agent réducteur.

16. Procédé selon la revendication 16, dens lequel la réaction est réalisée au voisinage de la neutralité. 17. Procédé selon la revendication 16, dans lequel l'agent réducteur est du cyanoborohydrure de

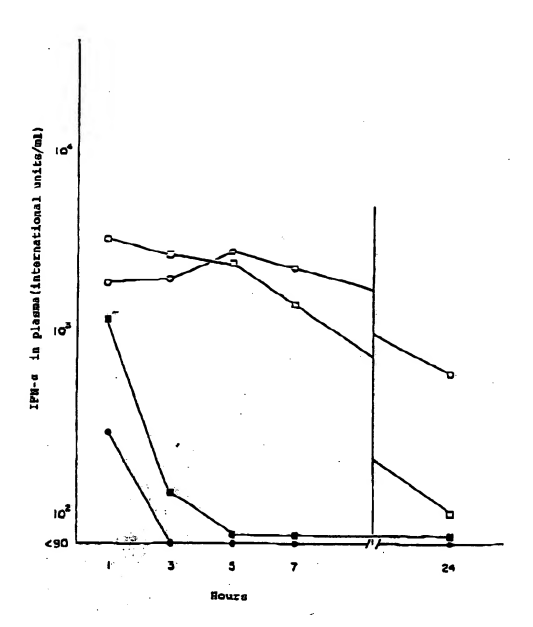
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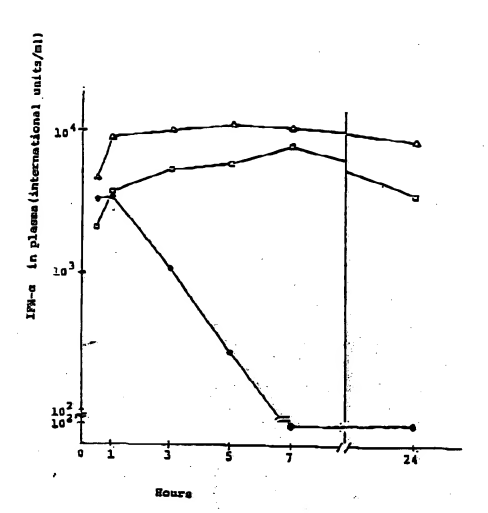
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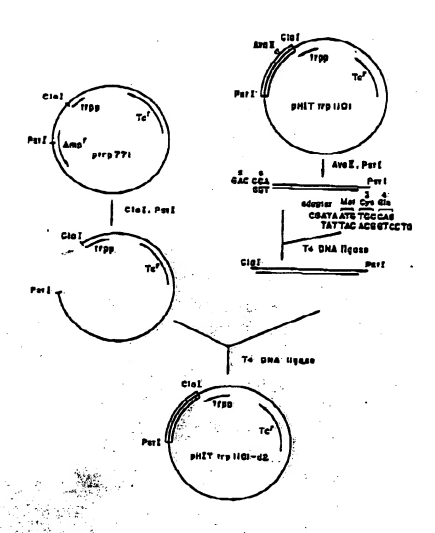
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F1g. 1





Pig. 3



Pig. 4

